LISTING OF THE CLAIMS

1. A biofunctionalized quantum dot, comprising:

a nanocrystalline core exhibiting quantum confinement and having a band gap and a surface;

a mercaptoalkanoic acid linked to the surface; and

a biofunctional group linked to the surface,

wherein the biofunctional group comprises a saccharide or the mercaptoalkanoic acid is linked to the surface of the nanocrystalline core without a shell layer.

2. The biofunctionalized quantum dot of claim 1,

the mercaptoalkanoic acid having exactly one carboxyl group and comprising less than seven carbon atoms.

- 3. The biofunctionalized quantum dot of claim 1, the mercaptoalkanoic acid comprising mercaptoacetic acid.
- 4. The biofunctionalized quantum dot of claim 1, further comprising: a shell layer overcoating the nanocrystalline core.
- 5. The biofunctionalized quantum dot of claim 4, the shell layer comprising cadmium sulfide or mercury sulfide; and the nanocrystalline core comprising cadmium telluride or cadmium selenide or mercury telluride or mercury selenide.
- 6. The biofunctionalized quantum dot of claim 1, the saccharide not comprising mannose or dextran.
- 7. The biofunctionalized quantum dot of claim 1,
 the saccharide being selected from the group consisting of a tumor-associated
 antigen and Thomsen-Friedenreich disaccharide.

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8. The biofunctionalized quantum dot of claim 1, the saccharide linked to a sulfur atom; and the sulfur atom linked to the surface of the nanocrystalline core.

- 9. The biofunctionalized quantum dot of claim 1, the saccharide linked to a linking group; the linking group linked to a sulfur atom; and the sulfur atom linked to the surface of the nanocrystalline core.
- 10. The biofunctionalized quantum dot of claim 9, the linking group comprising a carbon atom.
- 11. The biofunctionalized quantum dot of claim 1, wherein the biofunctionalized quantum dot is stable in aqueous solution under storage in the dark at 4 °C for at least 4 months with respect to luminescence, precipitation, flocculation, and leaching of the biofunctional group.
- 12. A formulation comprising the biofunctionalized quantum dot of claim 1 and further comprising a liquid,

wherein the biofunctionalized quantum dot is dissolved or suspended in the liquid and

wherein the biofunctionalized quantum dot does not precipitate or flocculate.

- 13. The quantum dot of claim 1, wherein the quantum dot comprises a therapeutic agent.
- 14. The quantum dot of claim 1, wherein the nanocrystalline core comprises a therapeutic agent or the biofunctionalized quantum dot further comprises a shell layer which comprises a therapeutic agent.

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15. A biofunctionalized quantum dot coated device, comprising: a device adapted for contact with a biological material and having a device surface; and

biofunctionalized quantum dots according to claim 1, wherein the biofunctionalized quantum dots are linked to the device surface to form a coating on the device.

- 16. A cell-quantum dot complex, comprising:
 the biofunctionalized quantum dot of claim 1;
 and a cell,
 wherein the biofunctional group is linked to the cell.
- 17. A method for producing a biofunctionalized quantum dot, comprising the steps of: providing a biofunctional group-thiol of Formula III; and,

Biofunctional Group
$$R_1$$
 SH III

refluxing the biofunctional group-thiol of Formula III with a cadmium salt, a hydrogen-alkali-group VIA element, and a suitable solvent to produce a quantum dot in a solution, wherein

 R_1 comprises a carbon atom and

the group VIA element is selected from the group consisting of tellurium and selenium.

- 18. The method of claim 17, the suitable solvent comprising water or N,N-dimethylformamide.
- 19. The method of claim 17, further comprising the steps of: reacting a glycoside of Formula I with an alkylthio acid in the presence of a

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catalyst to produce a thioester of Formula II;

Acetylated, Benzylidenated Biofunctional Group
$$R_1$$
 R_2

debenzylidenating the thioester of Formula II; and

hydrolyzing the thioester of Formula II to produce the biofunctional group-thiol of Formula III,

wherein R₁ comprises a carbon atom and R₂ comprises a carbon atom.

- The method of claim 17,the refluxing further comprising refluxing with a mercaptoalkanoic acid.
- 21. The method of claim 17, wherein the biofunctional group is a saccharide.
- 22. A method according to claim 17, further comprising the steps of:
 reacting a glycoside of Formula IV with an alkylthio acid in the presence of 2,2'azobisisobutyronitrile in 1,4-dioxane at about 75 °C to produce a thioester of Formula V;

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debenzylidinating the thioester of Formula V;

hydrolyzing the debenzylidinated thioester of Formula V to produce a Thomsen-Friedenreich-thiol of Formula VI; and

refluxing the Thomsen-Friedenreich-thiol of Formula VI with cadmium perchlorate, mercaptoacetic acid, hydrogen sodium telluride, and a suitable solvent,

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selected from the group consisting of water and N,N-dimethylformamide, to produce a Thomsen-Friedenreich-functionalized quantum dot in a solution.

23. A method of imaging, comprising the steps of: providing a biofunctionalized quantum dot according to claim 1; contacting the biofunctionalized quantum dot with a biological material; exposing the biological material to light having a wavelength effective to cause the quantum dot to fluoresce; and imaging the fluorescing quantum dots.

- 24. The method of claim 23, further comprising the step of using the imaging to identify tissue to which the biofunctional group exhibits high affinity as tissue in a diseased or abnormal state.
- 25. The method of claim 24, the diseased or abnormal state being cancerous.
- 26. A method of medical imaging, comprising the steps of: providing two types of biofunctionalized quantum dots according to claim 1, each type having a characteristic wavelength distinct from the other types;

each type of quantum dot functionalized with a different antigen or a different set of antigens;

contacting the two types of biofunctionalized quantum dots with a biological material;

exposing the biological material to light having a wavelength effective to cause the quantum dots to fluoresce; and

imaging the fluorescing quantum dots.

27. A method of therapy, comprising the steps of:

providing a biofunctionalized quantum dot according to claim 1; and

contacting the biofunctionalized quantum dot with a biological material and
thereby treating a disease.

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28. The method of claim 27, further comprising exposing the biological material to light having a wavelength effective to cause the quantum dot to fluoresce; and imaging the fluorescing quantum dot.

29. The method of claim 27, wherein the biofunctional group is selected from an immune-response stimulating group, a tumor-associated antigen, a Thomsen-Friedenreich disaccharide, and any combination of these.

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